

L9 ANSWER 7 OF 11 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.  
ACCESSION NUMBER: 1999:99102 BIOSIS  
DOCUMENT NUMBER: PREV199900099102  
TITLE: Inhibition of cell cycle progression by **rapamycin**  
induces **T** cell clonal **anergy** even in  
the presence of costimulation.  
AUTHOR(S): Powell, J. D.; Lerner, C. G.; Schwartz, R. H.  
CORPORATE SOURCE: LCMI, NIAID, NIH, Bethesda, MD USA  
SOURCE: Blood, (Nov. 15, 1998) Vol. 92, No. 10 SUPPL. 1 PART 1-2,  
pp. 21A.  
Meeting Info.: 40th Annual Meeting of the American Society  
of Hematology Miami Beach, Florida, USA December 4-8, 1998  
The American Society of Hematology  
. ISSN: 0006-4971.  
DOCUMENT TYPE: Conference  
LANGUAGE: English

L9 ANSWER 6 OF 11 CAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1998:300458 CAPLUS

DOCUMENT NUMBER: 128:320568

TITLE: Methods and materials for the induction of T cell **anergy**

INVENTOR(S): De Boer, Mark; Conroy, Leah B.

PATENT ASSIGNEE(S): Chiron Corp., USA

SOURCE: U.S., 35 pp., Cont.-in-part of U.S. Ser. No. 15,147.

CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 7

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5747034	A	19980505	US 1994-200716	19940218
US 5397703	A	19950314	US 1992-910222	19920709
US 5869050	A	19990209	US 1993-15147	19930209
CA 2183680	AA	19950824	CA 1995-2183680	19950119
WO 9522619	A1	19950824	WO 1995-US897	19950119

W: AU, CA, JP

RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

AU 9516877 A1 19950904 AU 1995-16877 19950119

EP 745136 A1 19961204 EP 1995-908634 19950119

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT,

SE

JP 09510607 T2 19971028 JP 1995-521804 19950119

PRIORITY APPLN. INFO.:

US 1992-910222 A2 19920709

US 1993-15147 A2 19930209

US 1994-200716 A 19940218

WO 1995-US897 W 19950119

AB Anti-B7-1 antibodies or other B7-1 ligands may be used to prevent or treat

a T-cell-mediated immune system disease in a patient or to induce antigen-specific tolerance. The anti-B7-1 antibodies may be used to

cause

T cell **anergy**, treat allograft transplant rejection, treat graft vs. host disease, and prevent or treat rheumatoid arthritis. An immunosuppressive agent is co-administered with the antibody.

L9 ANSWER 5 OF 11

MEDLINE

DUPLICATE 3

ACCESSION NUMBER: 1999172213 MEDLINE

DOCUMENT NUMBER: 99172213 PubMed ID: 10072524

TITLE: Inhibition of cell cycle progression by **rapamycin** induces **T** cell clonal **anergy** even in the presence of costimulation.

AUTHOR: Powell J D; Lerner C G; Schwartz R H

CORPORATE SOURCE: Laboratory of Cellular and Molecular Immunology, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, MD 20892, USA.

SOURCE: JOURNAL OF IMMUNOLOGY, (1999 Mar 1) 162 (5) 2775-84.  
Journal code: 2985117R. ISSN: 0022-1767.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 199904

ENTRY DATE: Entered STN: 19990426

Last Updated on STN: 19990426

Entered Medline: 19990414

AB Costimulation (signal 2) has been proposed to inhibit the induction of **T** cell clonal **anergy** by either directly antagonizing negative signals arising from TCR engagement (signal 1) or by synergizing with signal 1 to produce IL-2, which in turn leads to proliferation and dilution of negative regulatory factors. To better define the cellular events that lead to the induction of anergy, we used the

immunosuppressive

agent **rapamycin**, which blocks T cell proliferation in late G1 phase but does not affect costimulation-dependent IL-2 production. Our data demonstrate that full T cell activation (signal 1 plus 2) in the presence of **rapamycin** results in profound **T** cell **anergy**, despite the fact that these cells produce copious amounts of IL-2. Similar to conventional anergy (induction by signal 1 alone),

the

**rapamycin**-induced anergic cells show a decrease in mitogen-activated protein kinase activation, and these cells can be rescued by culture in IL-2. Interestingly, the **rapamycin**-induced anergic cells display a more profound block in IL-3 and IFN-gamma production upon rechallenge. Finally, in contrast to **rapamycin**, full T cell activation in the presence of hydroxyurea (which inhibits the cell cycle in early S phase) did not result in anergy. These data suggest that it is neither the direct effect of costimulation nor the subsequent

T

cell proliferation that prevents anergy induction, but rather the biochemical events that occur upon progression through the cell cycle

from

G1 into S phase.

L9 ANSWER 3 OF 11 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 2001:312008 BIOSIS

DOCUMENT NUMBER: PREV200100312008

TITLE: **Rapamycin** induces long term bone marrow chimerism in the absence of long term immunosuppression in

mismatched

stem cell transplantation; application to sickle cell

mice.

AUTHOR(S): Powell, J. D. (1); Fitzhugh, C. A.; Kang, E. M.; Weiss, S.;

Schwartz, R. H. (1); Tisdale, J. F.

CORPORATE SOURCE: (1) NIAID, NIH, Bethesda, MD USA

SOURCE: Blood, (November 16, 2000) Vol. 96, No. 11 Part 1, pp. 580a-581a. print.

Meeting Info.: 42nd Annual Meeting of the American Society of Hematology San Francisco, California, USA December 01-05, 2000 American Society of Hematology . ISSN: 0006-4971.

DOCUMENT TYPE: Conference

LANGUAGE: English

SUMMARY LANGUAGE: English

AB Engagement of the TCR leads to not only T cell activation but also upregulation of negative regulatory factors which promote tolerance. We have previously demonstrated that in contrast to cyclosporine (CSA) which inhibits anergy induction, **rapamycin** (Rapa) can promote T cell **anergy** even in the presence of costimulation. As such, we sought to determine if Rapa could be utilized to promote bone marrow chimerism in a Fl into parent transplantation model using minimal conditioning. Splenocytes (100 X 106) from G-CSF mobilized (C57BL/6 X BALB/C)Fl mice were injected into C57BL/6 recipients which had received 300cGy conditioning and either no immunosuppression (No IS, n=5), CSA (20mg/kg/d, n=6) I.P., or Rapa (3mg/kg/d, n=7) I.P. for 28 days. The No

IS

mice rejected their grafts by 1 week, while the CSA mice initially demonstrated donor-chimerism (10-15%) but eventually rejected their grafts. In contrast, the Rapa mice demonstrated progressively increasing donor chimerism which plateaued at 60-80%. More importantly, the Rapa

mice

have remained chimeric at this level for >3 months after stopping immunosuppression. Donor chimerism was preserved among CD4+, CD8+, B cell and granulocyte compartments. Donor cells were also detected in the thymuses of chimeric mice indicating a potential role for thymic

deletion.

In MLRs, splenocytes from both the No IS and CSA mice responded to BALB/c stimulator cells while cells from the Rapa mice were unresponsive. Using the Rapa protocol, mice thalassemic for murine Hb and transgenic for

human

HbS (expressing 60% human HbS) were transplanted with stem cells from normal Fl mice. Donor myeloid chimerism as low as 30% resulted in undetectable levels of HbS by Hb electrophoresis. Whether the virtually undetectable levels of HbS in the transplanted mice reflects a survival advantage of the normal RBC's or an advantage at the level of erythropoiesis is currently being investigated. In in vitro functional assays, blood from transplanted HbS trait mice displayed decreased turbidity in the sickle prep test as well as decreased or absent sickling on Sodium Bisulfite prepared smears compared to nontransplant controls. Current experiments to further characterize the rheologic properties of

chimeric mice as well as pathology in transplanted homozygous sickle mice are underway. Finally, these data suggest that this simple, non toxic, pharmacologic protocol might be useful in attaining hematopoietic chimerism in human allogeneic stem cell transplantation.

L9 ANSWER 2 OF 11 MEDLINE DUPLICATE 2  
 ACCESSION NUMBER: 2001106121 MEDLINE  
 DOCUMENT NUMBER: 20571934 PubMed ID: 11123330  
 TITLE: Relative resistance in the development of T cell  
 anergy in CD4+ T cells from simian immunodeficiency  
 virus disease-resistant sooty mangabeys.  
 AUTHOR: Bostik P; Mayne A E; Villinger F; Greenberg K P; Powell J  
 D; Ansari A A  
 CORPORATE SOURCE: Department of Pathology and Laboratory Medicine, Emory  
 University School of Medicine, Atlanta, GA 30322, USA..  
 pbstik@emory.edu  
 CONTRACT NUMBER: RO1 AI27057 (NIAID)  
 SOURCE: JOURNAL OF IMMUNOLOGY, (2001 Jan 1) 166 (1) 506-16.  
 Journal code: 2985117R. ISSN: 0022-1767.  
 PUB. COUNTRY: United States  
 DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
 LANGUAGE: English  
 FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals  
 ENTRY MONTH: 200102  
 ENTRY DATE: Entered STN: 20010322  
 Last Updated on STN: 20010322  
 Entered Medline: 20010208

AB Despite high viral loads, T cells from sooty mangabey (SM) monkeys that  
 are naturally infected with SIV but remain clinically asymptomatic,  
 proliferate and demonstrate normal Ag-specific memory recall CD4(+) T  
 cell  
 responses. In contrast, CD4(+) T cells from rhesus macaques (RM)  
 experimentally infected with SIV lose Ag-specific memory recall responses  
 and develop immunological anergy. To elucidate the mechanisms for these  
 distinct outcomes of lentiviral infection, highly enriched alloreactive  
 CD4(+) T cells from humans, RM, and SM were anergized by TCR-only  
 stimulation (signal 1 alone) and subsequently challenged with  
 anti-CD3/anti-CD28 Abs (signals 1 + 2). Whereas alloreactive CD4(+)T  
 cells  
 from humans and RM became anergized, surprisingly, CD4(+) T cells from SM  
 showed marked proliferation and IL-2 synthesis after restimulation. This  
 resistance to undergo anergy was not secondary to a global deficiency in  
 anergy induction of CD4(+) T cells from SM since incubation of CD4(+) T  
 cells with anti-CD3 alone in the presence of **rapamycin** readily  
 induced anergy in these cells. The resistance to undergo anergy was  
 reasoned to be due to the ability of CD4(+) T cells from SM to synthesize  
 IL-2 when incubated with anti-CD3 alone. Analysis of phosphorylated  
 kinases involved in T cell activation showed that the activation of  
 CD4(+)  
 T cells by signal 1 in SM elicited a pattern of response that required  
 both signals 1 + 2 in humans and RM. This function of CD4(+) T cells from  
 SM may contribute to the resistance of this species to SIV-induced  
 disease.

L82 ANSWER 4 OF 4 PCTFULL COPYRIGHT 2003 Univentio  
 ACCESSION NUMBER: 2000000825 PCTFULL ED 20020515  
 TITLE (ENGLISH): DETECTION AND MODULATION OF CELLULAR IMMUNITY TO  
 IMMUNE PRIVILEGED ANTIGENS  
 TITLE (FRENCH): PROCEDE ET AGENTS POUR LA DETECTION ET LA MODULATION  
 D'IMMUNITE CELLULAIRE SUR DES ANTIGENES PRIVILEGIES  
 IMMUNS  
 INVENTOR(S): DARNELL, Robert, B.; ALBERT, Matthew, L.; BHARDWAJ,  
 Nina  
 PATENT ASSIGNEE(S): THE ROCKEFELLER UNIVERSITY  
 LANGUAGE OF PUBL.: English  
 DOCUMENT TYPE: Patent  
 PATENT INFORMATION:

	NUMBER	KIND	DATE
	WO 2000000825	A2	20000106
DESIGNATED STATES	AU CA JP AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE		
APPLICATION INFO.:	WO 1999-US14827	A	19990630
PRIORITY INFO.:	US 1998-09/107,978		19980630
	US 1999-09/107,978		19990629

DETD In the practice of the above method, certain immune-privileged antigens  
 may not be adequately  
 taken up by **dendritic** cells for presentation on the  
 cell surface, nor will exposure of the  
 deDdritic cells to the intact antigen or its peptides. . . be  
 readily  
 processed and  
 presented. Among various known means for increasing antigen  
 presentation  
 by poorly  
 immunogenic or poorly processed antigens, use of **apoptotic**  
**cells** expressing the desired  
 antigen to deliver antigen to dendritic cells (17), in addition to  
 other  
 known means such as the  
 use of. . .